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Dated: May 20, 2008.

Gary D. Cooper,
District Manager.

[FR Doc. E8-11722 Filed 5-23-08; 8:45 am]

BILLING CODE 4310-GG-P

DEPARTMENT OF THE INTERIOR

Minerals Management Service

Extension of Post-Sale Evaluation Period for Central Gulf of Mexico Lease Sale 206

AGENCY: Minerals Management Service (MMS), Interior.

ACTION: Notice to Extend Post-Sale Evaluation Period for Central Gulf of Mexico Lease Sale 206.

SUMMARY: This notice extends by 30 days, the post-sale evaluation period for Central Gulf of Mexico Lease Sale 206. The Minerals Management Service (MMS) will complete evaluating all the bids received in this sale by July 17, 2008. This action is necessary due to the unusually high number of bids received in this lease sale.

DATES: The post-sale evaluation period ends on June 17, 2008.

FOR FURTHER INFORMATION CONTACT: David Marin, Regional Supervisor, Resource Evaluation, Gulf of Mexico Region, telephone 504-736-2710.

SUPPLEMENTARY INFORMATION: In the Central Gulf of Mexico Sale 206, held March 19, 2008, we received 1057 bids on 615 tracts, 513 tracts of which passed to a second phase requiring additional detailed evaluations. The aggressive

bidding activity is due, in part, to the high number of quality prospects on recently expired unexplored tracts in newly established deepwater hydrocarbon plays and to the cost saving technological advances related to hydrocarbon exploration and development in the Gulf of Mexico's deepwater environment. The unusually high number of bids received on a large number of tracts, and the high volume of exclusively reprocessed data identified on Sale 206, significantly increases the workload for reviewing the adequacy of bids. Consequently, MMS is unable to conduct and complete the bid review process within the 90 days, *i.e.*, by June 17, 2008. Under the provisions of § 256.47(e)(2), MMS is extending the bid evaluation period until July 17, 2008.

Dated: May 5, 2008.

Lars Herbst,

Regional Director, Gulf of Mexico OCS Region.

[FR Doc. E8-11711 Filed 5-23-08; 8:45 am]

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DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Manufacturer of Controlled Substances; Notice of Application

Pursuant to § 1301.33(a) of Title 21 of the Code of Federal Regulations (CFR), this is notice that on April 25, 2008, Siegfried (USA), Inc., Industrial Park Road, Pennsville, New Jersey 08070, made application by letter to the Drug Enforcement Administration (DEA) to be registered as a bulk manufacturer of Oripavine(9330), a basic class of controlled substance listed in schedule II.

The company will use the above listed controlled substance in the manufacture of other controlled

substance intermediates for sale to its customers.

Any other such applicant and any person who is presently registered with DEA to manufacture such a substance may file comments or objections to the issuance of the proposed registration pursuant to 21 CFR 1301.33(a).

Any such written comments or objections being sent via regular mail should be addressed, in quintuplicate, to the Drug Enforcement Administration, Office of Diversion Control, Federal Register Representative (ODL), Washington, DC 20537, or any being sent via express mail should be sent to Drug Enforcement Administration, Office of Diversion Control, Federal Register Representative (ODL), 8701 Morrisette Drive, Springfield, Virginia 22152; and must be filed no later than July 28, 2008.

Dated: May 15, 2008.

Joseph T. Rannazzisi,

Deputy Assistant Administrator, Office of Diversion Control, Drug Enforcement Administration.

[FR Doc. E8-11631 Filed 5-23-08; 8:45 am]

BILLING CODE 4410-09-P

DEPARTMENT OF JUSTICE

Drug Enforcement Administration

Manufacturer of Controlled Substances; Notice of Application

Pursuant to § 1301.33(a) of Title 21 of the Code of Federal Regulations (CFR), this is notice that on September 21, 2007, AMRI Rensselaer, Inc., 33 Riverside Avenue, Rensselaer, New York 12144, made application by renewal to the Drug Enforcement Administration (DEA) as a bulk manufacturer of the basic classes of controlled substances listed in schedules I and II:

| Drug | Schedule |
|---|----------|
| Marihuana (7360) | I |
| Tetrahydrocannabinols (7370) | I |
| Amphetamine (1100) | II |
| Lisdexamfetamine (1205) | II |
| Methylphenidate (1724) | II |
| Pentobarbital (2270) | II |
| Hydrocodone (9193) | II |
| Meperidine (9230) | II |
| Dextropropoxyphene, bulk (non-dosage form) (9273) | II |
| Oxymorphone (9652) | II |
| Fentanyl (9801) | II |

The company plans to manufacture bulk controlled substances for use in product development and for

distribution to its customers. In reference to drug code 7360 (Marihuana), the company plans to bulk

manufacture cannabidiol as a synthetic intermediate. This controlled substance will be further synthesized to bulk